CASE REPORT

FAMILIAL ACTINIC LICHEN PLANUS:
CASE REPORTS IN TWO BROTHERS

Shahriar Sadr-Ashkevari MD

Department of Dermatology, Razi Medical Center, Rasht, Iran

Abstract

We report two cases of a 17- and 19-year-old brothers from Rasht (North of Iran), who presented with actinic lichen planus lesions. The younger brother had typical annular patches on his forehead and neck and violaceous papules on his neck. The second brother presented with pigmented melasma-like lesions on his face. Histopathologic studies revealed lichen planus-like eruptions. Many reports describe the tendency of familial lichen planus (LP) developing at an early age and to becoming severe or chronic, and having widespread atypical manifestations. Our report probably is the first, which describes familial occurrence of actinic LP and adds further evidence to the possible role of genetic factors in triggering and determining different types of LP other than the disease itself.

Keywords • Lichen planus • familial • actinic • Iran

Introduction

Lichen planus (LP) is a unique papulosquamous dermatosis which may present with mucocutaneous lesions. It shows wide variation both clinically and histopathologically.

Factors including the consumption of various drugs or contact with heavy metals, systemic diseases, graft-versus-host disease, autoimmune liver disease, ulcerative colitis and myasthenia gravis have been implicated in the etiology of LP. These data suggest concurrent immunologic triggering and genetic susceptibility in the pathogenesis of LP.

LP has been reported worldwide, but is more prevalent among the younger age groups in both tropical and subtropical countries, with a predominance of some clinical types such as actinic LP that has been reported mostly in the Middle East and Asia.

In addition, LP has been reported in families and monozygotic twins, with an incidence of 1% to 11% of all LP cases. It has been characterized by its tendency to develop at an early age, become severe or chronic, and to have widespread atypical manifestations.

Case Presentation

Two Iranian brothers from northern Iran were referred to the Dermatology Department of Razi Medical Center in Rasht with cutaneous eruptions, both clinically and histologically compatible with actinic LP, in the summer of 1997.

Case 1

A 19-year-old male patient presented with brownish erythematous melasma-like patches, 2 x 3.5-4 cm in size, on his temporal and malar regions (Figure 1). The lesions had been present since 2 years prior to admission, but had shown aggravation during the summer with mild pruritus. The patient had not used any medications and no systemic disease was evident. Physical examination revealed no signs of LP in the other parts of the body, including the oral mucosa, hair or nails.

Case 2

A 17-year-old male patient (brother of case 1) presented with well-defined brownish erythematous patches with mild central atrophy on his forehead, glabella, nose, upper lip and sides of neck. These lesions varied in size and some, especially those on the sides of the neck, were annular in shape. A few typical LP papules were
Figure 1. Melasma-like brownish erythematous patches on the face and temple (Case 1).

Figure 2. Brownish erythematous patches with mild annular configuration on the forehead, glabella and upper lip (Case 2).

Discussion

It is assumed that LP results from a delayed type of hypersensitivity reaction against an unknown new epidermal antigen that develops either alone or concurrent with other diseases. Infective, psychosomatic, metabolic, pharmacologic, autoimmune, and genetic factors have been implicated in the etiologic of this disease.1,2,4 Reports exist on the occurrence of LP in families and monozygotic twins. Higher incidence of HLA-DR1 is detected in such patients.3,5,6,12 The familial occurrence of LP is a well-recognized but rare event, with an incidence varying from 1% to 11% of all LP patients. In comparison with classic LP, familial LP is characterized by its early age of onset, atypical and widespread clinical presentation and its higher tendency to become severe and chronic.10-13 Sodaify, in 1978, reported one of the largest familial groups of LP, including a Jewish mother and five of her nine children in Shiraz, without increased incidence of HLA class I phenotypes.12 On the other hand, actinic LP is a distinct variant characterized by annular erythematous brownish patches or plaques. In a few cases isolated or confluent papules, hypermelanotic melasma-like patches are confined to exposed parts of the skin, particularly the face, or even a dermatitis that occurs predominantly in Oriental (especially Middle Eastern) young people.7-12 Histologic findings reveal an interface dermatitis, mostly in the form of mild vacuolar degeneration of the basal layer, perivascular lymphocytic infiltration of the mid-dermis and significant pigmentary incontinence. The incidence of actinic LP was reported to be 0.9% in our previous study which was conducted on 110 cases of LP.14 With regard to this report, and to the occurrence of actinic LP in an Egyptian boy who had been living in England all his life, it is proposed that
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these case reports are not only the first on familial occurrence of actinic LP in the world, but also reveal further evidence that genetic factors (especially HLA-DR1) may be involved, somehow specifically, in the occurrence of LP and even its clinical appearance,3-6,10,12,13 a matter that must be confirmed through further studies.

With regard to the clinical melasma-like presentation in one of our cases, we agree with Al-Fuzan who suggested the inclusion of pigmented actinic LP in the differential diagnosis of patients presenting with facial melanosis.8

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References


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Figure 3. Basal cell degeneration, mild to moderate perivascular infiltration of lymphocytes and significant pigmentary incontinence (Case 1).

Figure 4. Severe basal cell degeneration with pigmentary incontinence and typical band-like infiltration of lymphocytes close to epidermis (Case 2).